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# The Environment Recording Unit in coral skeletons – a synthesis of structural and chemical evidences for a biochemically driven, stepping-growth process in fibres

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**Abstract.** This paper gathers a series of structural and biochemical in situ characterizations carried out to improve our knowledge of the fine scale growth patterns of fibres in coral skeletons. The resulting data show a clear correspondence between the mineral subunits of fibres and the spatial distribution of organic macromolecules. New observations using atomic force microscope confirm the close relationship between mineral and organic phases at the nanometre scale.

Synthesis of these data results in a significant change in our concept of the mineralization process in coral skeletons. In contrast to the usual view of an aggregate of purely mineral units independently growing by simple chemical precipitation, coral fibres appear to be fully controlled structures. Their growth process is based on cyclic secretion of mineralizing compounds by the polyp basal ectoderm. These biochemical components of the coral fibres, in which sulfated acidic proteoglycans probably play a major role, are repeatedly produced (proteoglycans are those glycoproteins whose carbohydrate moieties consist of long unbranched chains of sulfated amino sugars). This results in a stepping growth mode of fibres and a layered global organization of coral skeletons.

Therefore, in contrast to the widely accepted geochemical interpretation, we propose a fibre growth model that places coral skeletons among the typical “matrix mediated” structures. The crystal-like fibres are built by superimposition of few micron-thick growth layers. A biomineralization cycle starts by the secretion of a mineralizing matrix and the final step is the crystallization phase, during which mineral material grows onto the organic framework. Thus, each growth layer is the actual Environment Recording Unit.

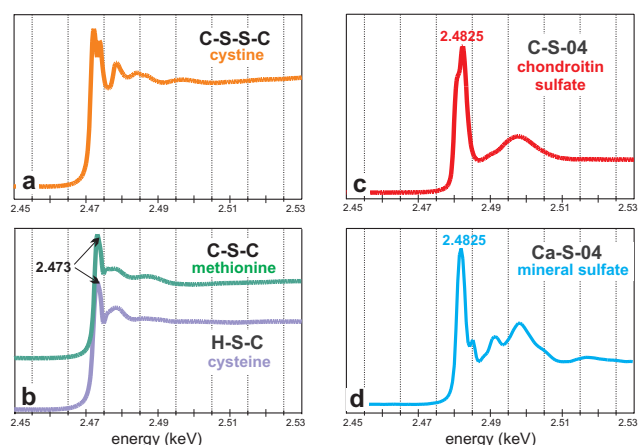
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From a practical standpoint, these results may contribute to develop a new high resolution approach of the environment recording by coral skeletons.

## 1 Introduction

Various chemical and/or isotopic proxies have placed coral skeletons among the most important sources of environmental information but, surprisingly, no agreement exists about the formation of these widely used biological archives. The process by which the basal ectoderm of coral polyps produces the underlying skeleton was first controversial in the late 19th century when, in contrast to the von Heider's theory (1881), it was recognized that calcification does not occur within the ectodermal cells themselves. Extracellular calcification occurring outside the ectodermal cell layer, as advocated by von Koch (1882), gained general agreement. Since then, the level of biological control over coral skeleton remains an unsolved question. As summarized by Le Tissier (1991), models of calcification in corals range from a pure physicochemical (Barnes, 1970) to a biologically controlled process (Johnston, 1977, 1980).

Crystallization of coral aragonite being an extracellular phenomenon, the hypothesis of a biological control exerted on skeleton formation should be supported first by data concerning the place where crystallization occurs: the interface between the polyp's basal cell layer and the underlying skeleton. In spite of Goreau's pioneering studies (1956–1959), research focussing on physico-chemical characteristics of the subectodermal space are extremely rare. In contrast to Molluscs (e.g. Pelecypods), where the mineralization space between the outer side of the mantle and the shell is more easily



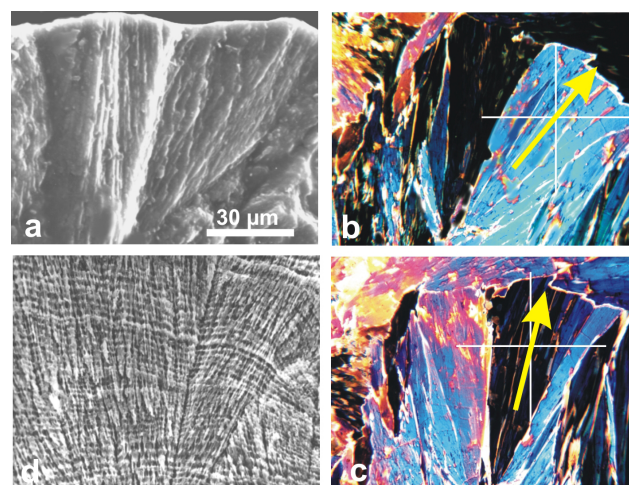
**Fig. 1.** Differences between XANES spectra from four sulfur organic bounds and sulfur in a calcium sulfate. Not only is the edge energy significant, but also the “near edge” oscillation: mineral and organic sulfates are clearly different.

accessible, the complexity of coral skeletons prevents easy access to the interface compartment. Extraction of mineralizing fluids to study their compositions and mineralizing capabilities was carried out in Molluscs as early as 1974–1976 (Wada and Fujinuki, 1974, 1976). To date, no equivalent research exists for corals. The first pH measurement at the interface between the basal ectoderm and the growing surface of a coral skeleton was only attempted in 2003 (Al-Horani et al., 2003).

Therefore, instead of a direct analysis of the mineralizing sites, complete decalcification of coral skeletons providing access to organic materials entrapped during skeletal growth has been much more used. This resulted in a series of papers emphasizing the importance of, either sugars (Wainwright, 1963; Wilfert and Peters, 1969; Dauphin, 2001), or amino acids (Young, 1971, 1973; Mitterer, 1978; Constantz and Weiner, 1988; Cuif et al., 1999). Nevertheless, as pointed out by Johnston (1980), research dealing with the biochemical composition of organic materials extracted from coral skeletons has resulted in mineralization models made “in ignorance of this material’s spatial distribution and micro-architecture within the skeleton”.

Moreover, additional uncertainty persists with respect to the amount of these organic components. Since Wainwright (1963), the fraction of organic material embedded in the coral skeletons has been commonly estimated at 0.1% of weight, a proportion recently re-evaluated to about 1% (Cohen and McConnaughey, 2003). Thermogravimetric measurements (Cuif et al., 1997) indicate that 2.5 to 3% of weight of coral skeletal carbonate was lost prior to the thermal decomposition of aragonite. Such discrepancies cast doubts on the actual influence of these organic compounds on the crystallization process.

In response to Johnston’s remark, this paper brings together a series of data based on a different approach. High



**Fig. 2.** Three different aspects of the same group of coral fibres. (a) Morphology of fibres on a fracture surface of a corallite (*Favia stelligera*). (b–c): Ultra-thin section observed in polarized light. Fibres are groups of subunits with overall similar behaviour with respect to polarization directions (orientation of the yellow arrow). (d) Aspect of the same skeletal sector after polishing and acidic etching. Overall organization of fibres is still recognizable, but growth layers have been made visible by synchronous differences in sensitivity to dissolution. The coordinated changes in thickness and spacement of growth layers become visible, allowing a precise description of fibre growth to be made.

resolution analytical instruments (scanning electron microscopy, synchrotron X-ray fluorescence and atomic force imaging) now allow in situ physico-chemical characterization of coral skeleton units at the micrometric and nanometric levels. Combining these data results in a new set of information about structure and composition of the skeletal units, the crystal-like fibre, and allows the reexamination of the calcification process itself.

## 2 Material and methods

Coral specimens have been collected alive, mostly by the authors during recent years. To remove tissues, polyps were decayed by a three-hour immersion in distilled water that causes the cells to be destroyed. Removal of the remaining living tissues was done by using a water-jet, followed by immersion in 0.1M sodium hypochlorite to clean the deeper parts of skeletons. Then, specimens were rinsed and air dried at room temperature. In all cases, only the uppermost parts of skeletons were used for imaging. Specimens were selected by SEM observation on polished and etched surfaces, which allow the fibrous microstructures to be examined. Samples contaminated by endolithic borers were discarded.

## 2.1 Materials

Corallites belonging to the following species have been used: *Favia stelligera* (Moorea Island, Polynesia); *Montastrea curta* (Mururoa Island, Polynesia; J. P. Chevalier coll.); *Diploastrea heliopora*, *Leptoria phrygia*, *Porites* cf. *australiensis*, *Acropora digitifera*, *Merulina scabricula* (New Caledonia lagoon); *Caryophyllia smithi* from Mediterranean Sea; *Cladocora caespitosa* (Mediterranean Sea, Marseille coastal region); *Favia fragum* (Guadalupe Island, Caribbean).

## 2.2 Methods

### 2.2.1 Structural and chemical characterizations

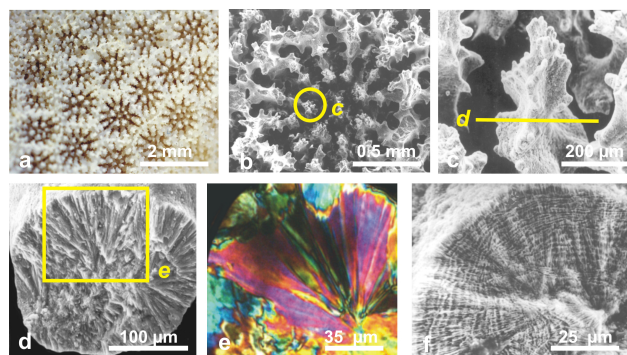
Microstructural features of fibres were first checked by microscopic observation of ultra-thin sections in polarized light. Ultra-thin sections, an improvement of the usual petrographic techniques, require fine polishing of the sample surface and a reduced thickness of slides. Instead of the standard 30 microns, grinding is continued to 4–5 micrometers thickness, allowing the primary colours of aragonite to be visible. Final polishing of the upper surface ensures an accurate microstructural observation in polarized light.

### 2.2.2 Scanning Electron Microscopy (SEM)

Observations were carried out not just on simple fractured surfaces but more often on polished and etched surfaces. Etching by very light acidic solutions (one per mil formic acid with addition of glutaraldehyde 3 to 5%) or enzymatic solutions reveals that differences in solubility do exist within the crystal-like fibres. Additionally, the overall organization of coral septa is well shown, by emphasizing the trace of the early mineralization zones (commonly called “centres of calcification”).

### 2.2.3 X-ray Absorption Near Edge Spectrum (XANES) characterization and mapping of sulfated polysaccharides

At the European Synchrotron Radiation Facility in Grenoble (France), the ID-21 instrument is a high resolution X-ray fluorescence detector. A high performance X-ray monochromator allows a very precise wavelength selection between 3 and 8 keV. This makes possible to select the energy bounds corresponding to different oxidation states of a given element. To characterize the sulfur state in coral skeletons, a calibration phase allows the different XANES spectra from sulfated standard molecules to be clearly separated (Fig. 1a to d). Sulfur in sulfated amino acids (methionine and cysteine), disulfur bonds in cystine and sulfated sulfur in chondroitin sulfate provides distinct spectra from both peak absorption wavelength and the XANES sectors of the absorption spectrum. This allows to characterize the state of organic sulfate in situ. High performance focussing lenses allow this



**Fig. 3.** From morphology to elemental growth layer in a *Porites* skeleton. (a–b) Morphology of *Porites* cf. *australiensis* corallite. (c) Longitudinal view of a vertical unit. (d) Radial disposition of fibres in a vertical unit. (e) Ultra-thin slide in a vertical unit: radial disposition of fibre fans is well visible. This is a typical example of what led numerous authors to emphasize the similarity of coral fibres with abiotic crystallisations. (f) Etching of the fractured surface in the *Porites* corallite, reveals the fibre growth layers.

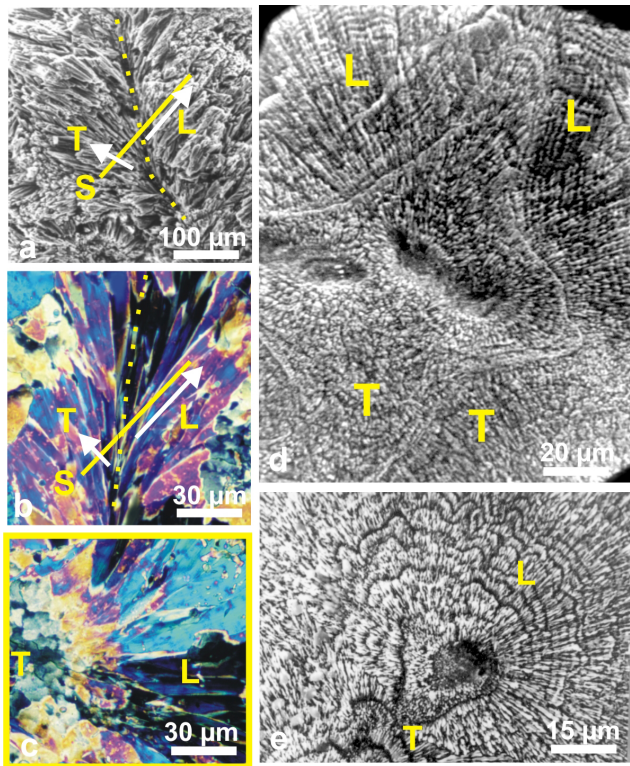
characterization to be obtained with sub-micron spatial resolution. A two-dimensional piezo-electrically driven specimen holder makes possible a micron-level displacement of the sample polished surface, resulting in a biochemical map of the skeletal structure.

### 2.2.4 Atomic Force Microscopy (AFM)

Scanning probe microscopy encompasses a family of techniques that measures surface topography and properties at an atomic scale. The atomic force microscope simultaneously produces maps the surface topography (height images or derived amplitude images) and phase images. In phase imaging, a variant of tapping mode, the phase lag of the cantilever oscillation relative to the signal sent to the cantilever's piezo driver is used as a basis for image generation. Phase images can be generated as a consequence of variations in material properties such as composition, visco elasticity, adhesion. AFM observations were conducted with Digital Instruments (Veeco) Nanoscope III Dimension 3100 at room temperature and air. The probe consisted of a cantilever with integrated Si<sub>3</sub>N<sub>4</sub> tips (Digital Instruments). Micron scale images were acquired using tapping mode.

For AFM observation of biominerals, there is presently no applicable routine procedure. Various preparatory processes have been used, all of which aiming to reduce the possible changes in relationship between mineral and anticipated organic components. Consistency between phase and height images is essential for interpretation. The procedures of the sample preparations are given in the figure captions.



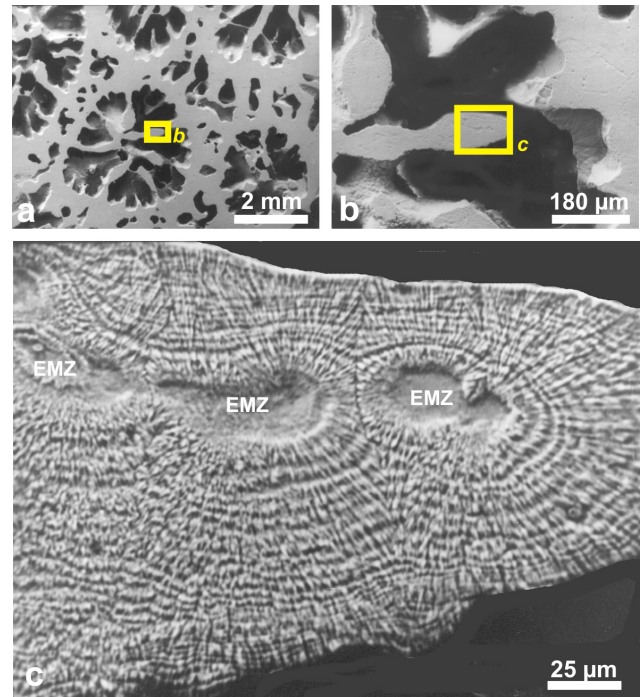


**Fig. 4.** Diversity of fibre directions on a corallite section, and consequences on the observation of etching results. (a) Symmetrically diverging fibres on both sides of a septum median plan (yellow dotted line). (b) Diverging fibres observed in thin slides, polarized light. Here the fibre fan-system is observed on a longitudinal (axial) section. The yellow line S corresponds to the perpendicular section (c, below). (c) This yellow framed view illustrates what is seen when the orientation of the section corresponds to the yellow line of (b). Depending on their orientation, fibres may appear transversely (T) or longitudinally (L) cut. On the left side, fibres appear as polygonal units because they are transversely cut (T on this picture and the S line of picture b). On the right side, fibres are longitudinally cut (L). (d–e) On the etched surfaces, fibre growth layers are well visible in longitudinal sections only: (d) *Leptoria phrygia*; (e) *Diploastrea heliophora*.

### 3 Results

#### 3.1 The incremental growth mode of fibres

SEM observation and ultrathin sections of coral fibres observed in polarized light allow to check their common characteristic: they appear as groups of elongated units (Fig. 2a) with a monocrystalline behaviour (Fig. 2b–c). However, etching of this polished surface reveals a completely different pattern. Differences in solubility result in a series of lines that demonstrate the incremental growth of fibres (Fig. 2d). This result is exemplified here by a *Porites* cf. *australien-sis* colony (Fig. 3a–b). The fractured skeletal rod (Fig. 3c) shows the radial organization of fibres (Fig. 3d). An ultra-



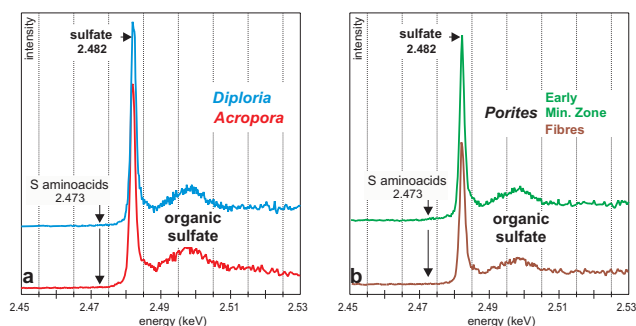
**Fig. 5.** Synchronism of growth layers in a septum of *Favia stelligera*. Growth of fibres is coordinated at an overall level that suggests a synchronism in mineralizing activity of the polyp ectodermal cell layer. EMZ=Early Mineralization Zone: at the distal tips of the microstructural units, calcification is initiated by specific structures that exhibit microstructural and compositional patterns (see also Fig. 7d).

thin section (Fig. 3e) confirms the monocrystalline behaviour of fibres, whereas the concentric growth lines are well visible after etching (Fig. 3f).

To date, no exception is known to this micron-scaled incremental growth of coral fibres. Attention must be drawn to the fact that the appearance of the stepping growth pattern depends on the orientation of the fibres with respect to the observation surface. It has long been recognized that tridimensional arrangements of fibres are complex and very different among corals from various families. With respect to growth direction (Fig. 4a, white arrows) the fibres are overall oblique to the septal median plan or growth axes. Consequently, within a given surface (Fig. 4b, line S), fibres may appear under different sections between two extreme conditions exemplified in Fig. 4c: from longitudinal (L) to transverse sections (T). Figures 4d and e show equivalent SEM pictures, in which appearance of the incremental growth pattern depends on the orientation of fibres.

#### 3.2 Global coordination of growth increments

Etched surfaces also show that high/weak solubility zones are continuous between adjacent fibres. Sometimes continuity of growth lines can be observed on the whole septum



**Fig. 6.** Responses of polished surfaces in corallites of *Diploria labyrinthica*, *Acropora digitiformis* and *Porites cf. australiensis* to a 2.4825 keV X-ray beam. In all cases a strong signal of organic sulfates is detected, whatever the location of the X-ray beam on the corallite surface, including the Early Mineralization Zones (the “centres of calcification”). In contrast, no response is obtained for S amino acids.

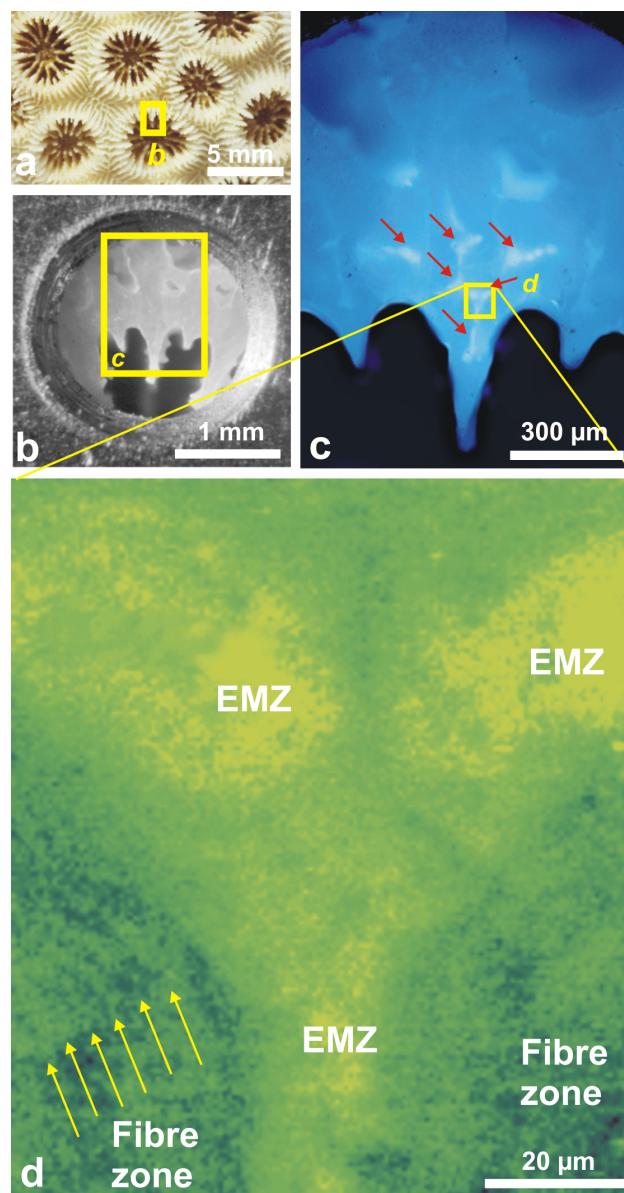
(Fig. 5a–c). Such a concentric growth pattern shows that thickening of septa is ensured by superimposition of growth layers that result from a global coordination of the biomineralization process on wide areas of the basal ectoderm. Additionally, Fig. 5c also shows that the series of superimposed growth layers surrounds regions that exhibit a distinct microstructural pattern. These regions correspond to the distal tips of septa, the regions where the calcification is initiated. These early mineralization zones (Fig. 5c: EMZ) are the long controverted domains called “centres of calcification” (Cuif and Dauphin, 1998).

### 3.3 XANES mapping: layered distribution of sulfate within fibres

All polished surfaces exposed to the 2.4825 keV X-ray energy produce a strong signal corresponding to the presence of organic sulfated polysaccharides (Fig. 6). In contrast, sulfated amino acids are practically undetectable. To take the full advantage of the ID-21 mapping device, a preliminary exploration of the surface is useful. Figures 7a to c show part of a polished section (wall and septa) in a *Montastrea* corallite exposed to UV light (365 nm) on a reflection microscope, allowing the EMZ zones to be localized (Fig. 7c: red arrows). The 2.4825 keV beam is then applied to this sector, the distance between measurement points and lines being one micron. Time exposure for each point is 0.8 s.

This results in the map of sulfate (Fig. 7d), between EMZ and fibres. Presently no precise quantification is possible, owing to the heterogeneity of the biogenic mineral that does not allows any calculation concerning the absorbed/emitted radiations. However, the higher concentration of sulfated sulfur in EMZ is well visible, as well as the banding patterns of sulfated sulfur in the fibrous part of the field view (arrows).

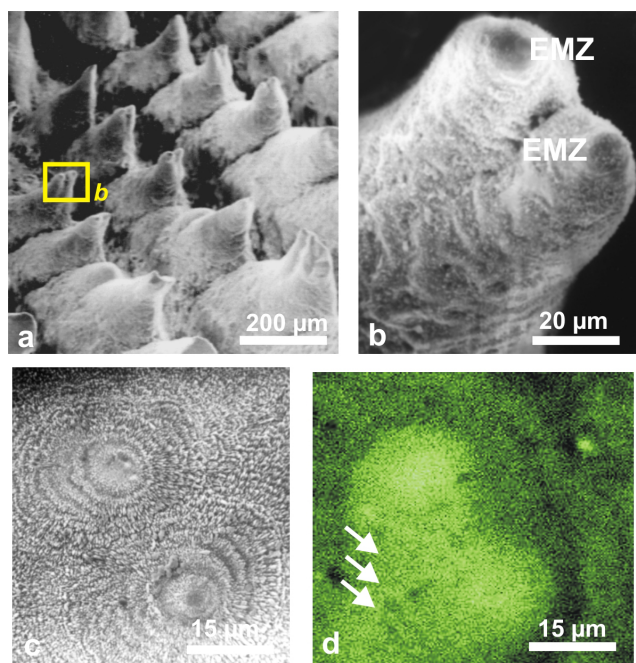
Very comparable results are obtained on corallites from other species: *Diploria labyrinthica* (Fig. 8a–d), *Porites cf.*



**Fig. 7.** Mapping of sulfated polysaccharides in *Montastrea curta*. (a) Morphology of the corallites. (b) Specimen in the sample holder. (c) Selection of the region to be mapped by UV fluorescence: the Early Mineralization zones show a strong response (red arrows). (d) Biochemical map of the selected zone. The Early Mineralization Zones exhibit a high sulfated polysaccharide concentration. In fibres, the well visible banding pattern in exact conformity with fibre growth layers (arrows) shows that mineral phase and sulfated polysaccharides are associated at a submicrometer scale within fibres.

*australiensis* (Fig. 9a–b), *Acropora digitifera* (Fig. 9c–d). In all samples, a correspondence is well visible between microstructural patterns and the distribution of sulfate.



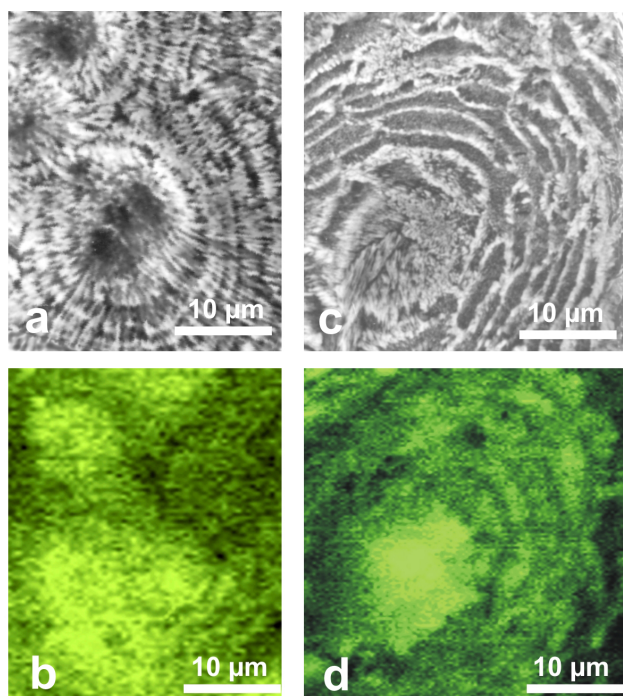


**Fig. 8.** Growth layers and sulfated polysaccharide distribution in *Diploastrea labyrinthica*. (a) Morphology of growing edges of septa built by a simple series of conical units. (b) At the tip of conical units, the Early Mineralization Zone (EMZ) surrounded by the beginning of fibrous zone. (c) Etched section, showing the concentric growth layers surrounding the EMZ. (d) XANES mapping of organic sulfur provides a very comparable pattern.

### 3.4 Heterogeneity of fibres at the nanometric scale

AFM examination confirms that fibres are not homogeneous structures at the submicrometer scale. Pictures of *Merulina scabricula* are particularly significant. Densely packed nanograins are well visible (Fig. 10a–b). Looking more closely (Fig. 10c–d), the grains seem coated by a thin layer (Fig. 10d). Due to the effectiveness of AFM phase imaging to detect physico-chemical properties, we can assess that the thin layer on the grain surfaces (Fig. 10c, arrows) is different from the grains themselves with respect to its chemical composition (Fig. 10d). Remarkably, similar pictures have been obtained on skeletons of every studied species, and at any place in the fibrous structures of each specimen (Fig. 11a–b: *Favia stelligera*; Fig. 11c–d: *Cladocora caespitosa*; Fig. 11e–f: *Caryophyllia smithi*).

These results show that: (1) a nanometric granular structure is the basic organization of the fibre growth layers and (2) the fibrous tissue is basically heterogeneous: the few tenth to hundred nanometre grains are included within a material that exhibits specific physico-chemical properties.



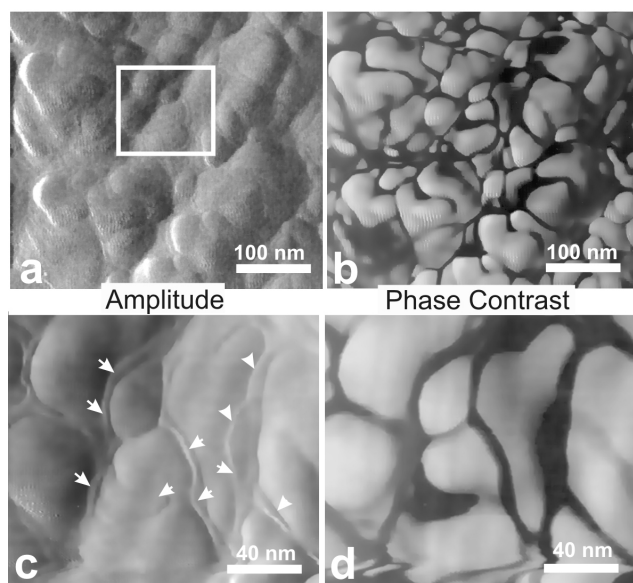
**Fig. 9.** Microstructural patterns and corresponding XANES mapping of organic sulfate on polished surfaces in *Porites* cf. *australiensis* (a–b) and *Acropora digitifera* (c–d).

## 4 Discussion

From McConeaughey (1989) to Sinclair and McCulloch (2004), geochemical interpretations have usually considered that coral aragonite is produced by a simple crystallization process occurring in a fluid of which composition is “close to sea water”. In this model, biological activity is limited to “enzymatic ion transport” of  $\text{Ca}^{2+}$  that creates the required conditions of supersaturation for  $\text{CaCO}_3$  precipitation. In this concept, no mention is made of the skeletal organic compounds. Their role in the crystallization process and their possible influence on Ca-carbonate composition due to their chemical activity are not taken into account. Clearly, as emphasized by Johnston (1980), absence of significant information about the distribution of the “organic matrix” has long remained a major obstacle to progress in understanding its importance.

### 4.1 Organic matrix at the nanoscale: consistency of AFM data with Johnston’s results

When Johnston published his transmission electron microscope pictures (1977, Fig. 1; 1980, Fig. 18), the organic network shown within the uppermost part of a coral fibre was the first evidence of a possible relationship between organic and mineral materials at a submicrometer scale. Of course, no mineral was visible within the network spaces, due to

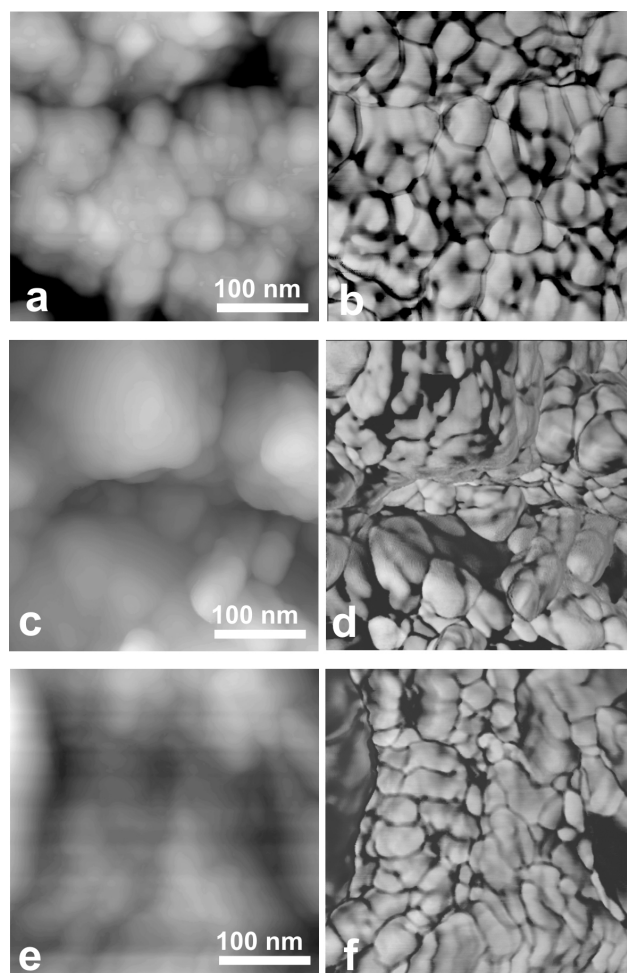


**Fig. 10.** AFM images of skeletal nanograins in *Merulina scabricula*. Amplitude images (a and c) and phase images (b and d) show the dual composition of skeletal grains. Phase imaging reveals the importance of the very weak relief that can be seen on (a) and (c) pictures of the grain surfaces (arrows). The very high contrast produced by these structures demonstrates that they are basically different, from a chemical standpoint. The XANES in situ characterization of sulfated polysaccharides at a submicron scale fully supports the interpretation of this high phase-contrast material as an organic coating of skeletal grains.

the preparation process used for transmission electron microscopy.

On the AFM pictures, dimensions of the nanograin's envelopes (Figs. 10–11) remarkably correspond to those of the organic network cells in the Johnston's pictures. This dimensional correspondence supports the hypothesis of an organic nature for the material separating nanograins within fibres, as already suggested by the interactive properties of this material in AFM phase imaging (in this technique, black means "attractive").

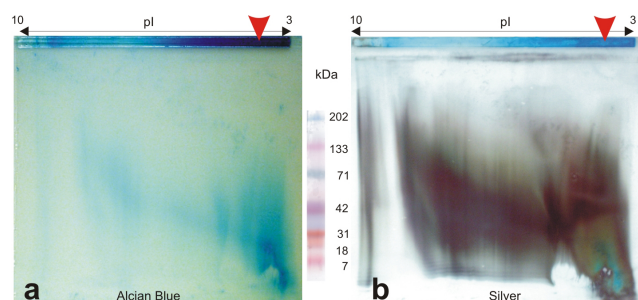
Owing to the used preparation process utilized, Johnston was able to characterize the organic network only in the uppermost  $3\ \mu\text{m}$ . at the top of the fibres. Therefore he suggested that the organic network might "disappear" in the deeper parts of the skeleton. Three different results contribute to disprove this suggestion. (1): no change is visible in AFM nanograin envelopes far deeper than Johnston thought, even at more than one centimeter below the septal growth surface (several hundreds of superimposed growth layers). (2): etching shows that the layered organization of skeletons remains visible all along the fibre length (e.g. Fig. 4). (3): the XANES signal indicating sulfated sulfur is obtained on the whole surface of coral sections, whatever the distance to the growing surface.



**Fig. 11.** Equivalent AFM amplitude and phase pictures from coral-lite belonging to three species: (a–b) *Favia stelligera*; (c–d) *Cladocora caespitosa*; (e–f) *Caryophyllia smithi*. Note that surface topography remains rather unprecise on height images (a, c, e), but phase imaging (b, d, f) provides clear information about the basic granular structure, that has been found in all coral skeletons observed to-date.

Particular attention should be paid to the XANES sulfated sulfur signal because sulfur in coral skeletons has long been reported (see Milliman, 1974). It was variously interpreted: as small amounts of  $\text{CaSO}_4$  (Milliman), as sulfate ions (Takano et al., 1980) or sulfite (Bar-Matthews, 1993) substituting for  $\text{CO}_3$  anions. The suggestion that S might be a component of mineralizing matrices have also been made (Blake and Peacor, 1981; Mackenzie et al., 1983). Applying the XANES method to a variety of inorganic and organic carbonates, Pingitore et al. (1995) noted the "specificity and exclusivity" of XANES spectra but, following the interpretation of Takano et al. (1980), they supported the mineralogical interpretation of  $\text{SO}_4$  ions being substituted for  $\text{CO}_3$ .

In contrast to these views inspired by a purely mineral concept of fibres, the presence of organic sulfate in coral



**Fig. 12.** 2-D electrophoretic characterization of sulfated polysaccharides and protein compounds from coral skeletons. Alcian blue staining of acidic sulfated polysaccharides shows that, after isoelectric focussing has established their low isoelectric point (pI), most of them remain included in the first dimension strip gel. This shows that their molecular weights exceed 300 kDa, the upper molecular weight accepted by the gel.

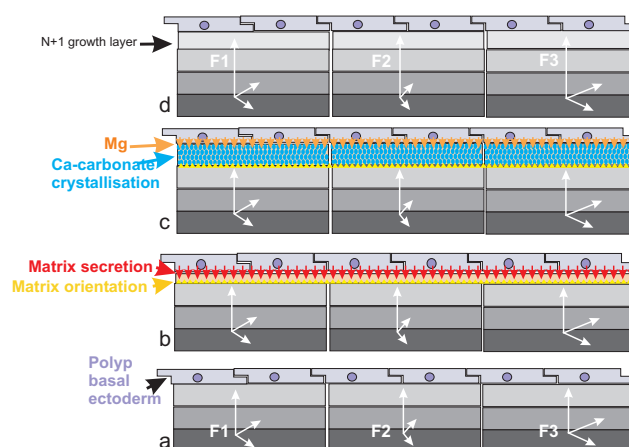
skeletons is now well established by Alcian Blue staining of 2-D electrophoresis gel (Dauphin, 2001). Not only the organic nature of sulfate is thus established, but the very high molecular weight of the sulfated polysaccharides is also demonstrated, as well as their highly acidic composition (Fig. 12). Additionally, in contrast to Pingitore's interpretation based on punctual XANES spectra only (without any microstructural data), XANES maps show a layered distribution of sulfated sulfur. These properties are very consistent with a possible role of sulfated polysaccharides in fibre mineralization.

#### 4.2 “The hypothesized organic matrix mediation of crystal growth (Johnston, 1977; Jackson and Gladfelter, 1985) is difficult to support”

This statement (Constantz, 1986, p. 155) illustrates Le Tissier's comment (Le Tissier, 1991) about the diversity of biomineralization models for coral fibres. A major obstacle to progress in this question was the difficulty to connect microstructural data showing the layered organization within coral fibres (and numerous other biominerals: Cuif et al., 1981, 1997) to chemical or biochemical data obtained at the corresponding scale.

By microprobe analysis, microstructure-linked differences in sulfur concentrations were shown in the septa of a number of coral species (Cuif and Dauphin, 1998), drawing attention to the specific chemical properties of the “centres of calcification”. Located at growing tips of septa, they initiate the mineralization process (early mineralization zones: EMZ). The XANES maps fully confirm this biochemical difference between the EMZ and the ectoderm zones that produce fibres.

Once the biochemical difference between EMZ and fibres is well established, attention must be focussed to the XANES maps that prove the equivalence between the distribution of



**Fig. 13.** Scheme of a growth layer formation in a coral skeleton, summarizing the structural, chemical and biochemical data. F1, F2, F3: upper mineralized layers in three adjacent fibres; arrows: overall crystallographic orientations.

organic sulfate and the mineral growth layers (Cuif et al., 2003). This correlation implies that, within each growth layer, organic components are associated with the mineral phase itself at a submicron scale. The AFM data discussed above (4.1), provide us with an explanation of this intriguing correlation between organic matrix and mineral layers. Actually, relationships between mineral and organic components have to be established and understood at the nanometre scale. Clode and Marshall (2002) have clearly shown a network of organic filaments that creates a link between the outer face of ectodermal cells and the upper surface of aragonite fibres. Their conclusion concerning the potential role of this network in the biomineralization process is fully supported par the AFM pictures.

From a quantitative stand point, recent results of ATG measurements coupled to infrared spectrometry (Cuif et al., 2004) have confirmed that 2.5–3% of the coral skeleton weight is lost before thermal decomposition of aragonite. Additionally, these measurements have shown the dual origin of these weight losses. About 0.8 to 1% by weight (depending upon species) is due to loss of organic material properly speaking, a result that supports the estimation presented by Cohen and McConnaughey (2003). The other part of the weight loss is due to water linked to organic molecules.

This hydrated nature of the skeletal organic material is also consistent with the previous biochemical characterization (Dauphin, 2001). Glucids in the fibre mineralizing compounds belong to the family of glycoconjugates well known for their ability to include water in their complex molecular assemblages. Therefore, the presence of water that has long been recognized in skeletons (Gaffey, 1988), is not surprising. Due to its submicroscopic distribution within the skeleton, Gaffey concluded that water was located as “minute inclusions”. Biochemical characterizations associated to ATG



analysis suggest that water is directly associated to sulfated polysaccharides at the molecular scale. From the same quantitative point of view, it must also be noted that a 2.5–3% ratio in weight for the hydrated organic compounds indicates that even assuming a high density for this hydrated organic phase (i.e. 1.5), we can estimate that about 5–7% of the skeleton volume is made of non-carbonate components (aragonite density: 2.99). As a first approach, this result also appears rather consistent with the spatial ratio between organic material and nanograins in AFM pictures.

#### 4.3 A polycyclic model for the growth of the coral fibres

The structural and chemical data reported here can be summarized by a simple model of the growth layer formation (Fig. 13).

At the starting point of a biomineralization cycle, the basal ectoderm of the polyp is in close contact with the upper surfaces of fibres (Fig. 13a: F1 to F3), each of them having its *c* axis in overall conformity with elongation of each fibre (note that orientation of *a* and *b* axes differs from fibre to fibre – see Fig. 2).

##### 4.3.1 Organic secretion and formation of the matrix framework

Although the physiology of matrix secretion is thoroughly studied (see Tambutté, 1996; Allemand, 2004 for review), molecular organization outside the ectodermal layer is still poorly understood. Chromatographic and electrophoretic characterizations of coral matrices have emphasized the importance of acidic, high molecular weight glucidic compounds (Dauphin, 2001; Dauphin and Cuif, 1997). The ability of molecules of this biochemical family to produce polymeric structures is known and relevant to their potential role in the first step of the biomineralization cycle. Radiometric methods have also shown the presence of low weight peptides (Puverel, 2004). This complex blend of high molecular weight sulfated sugars and acidic proteins with medium to low molecular weight suggests that within the mineralizing compartment (outside the ectodermal cells) a self-assembly process may occur, resulting in a chemically active organic framework. The historical observation by Goreau (1956) of a glucidic layer on the external side of the basal ectoderm of the polyp along with the network of organic filaments recently observed by Clode and Marshall (2002) using cryofracturing methods may correspond to this secretory/ polymerization step.

This self-assembly step may also be a relevant way to explain the overall crystallographic continuity of fibres. Cytological observations by transmission electron microscopy (Tambutté, 1997) have shown the complex arrangement of the ectodermal cells, “frequently overlying each other” (Clode and Marshall, 2003, Fig. 1). No correspondence exists between a given fibre and the cells of the basal ectoderm:

thus the crystallographic continuity of fibres cannot directly result from the organization of ectodermal cells. At the beginning of the biomineralization cycle, the macromolecular assemblage is secreted in close contact with the upper surface of aragonite fibres facing the basal ectoderm. Numerous observations (Mitterer and Cunningham, 1985; Hazen et al., 2001) have demonstrated the interactivity between Ca-carbonate crystal surfaces and organic molecules. Thus, the crystal lattice of fibres may act as a template during the first step of the self-assembly process, resulting in the orientation of the matrix organic framework in conformity with the crystallographic orientation of each of the underlying fibres (Fig. 13b).

##### 4.3.2 Crystallization within the matrix framework

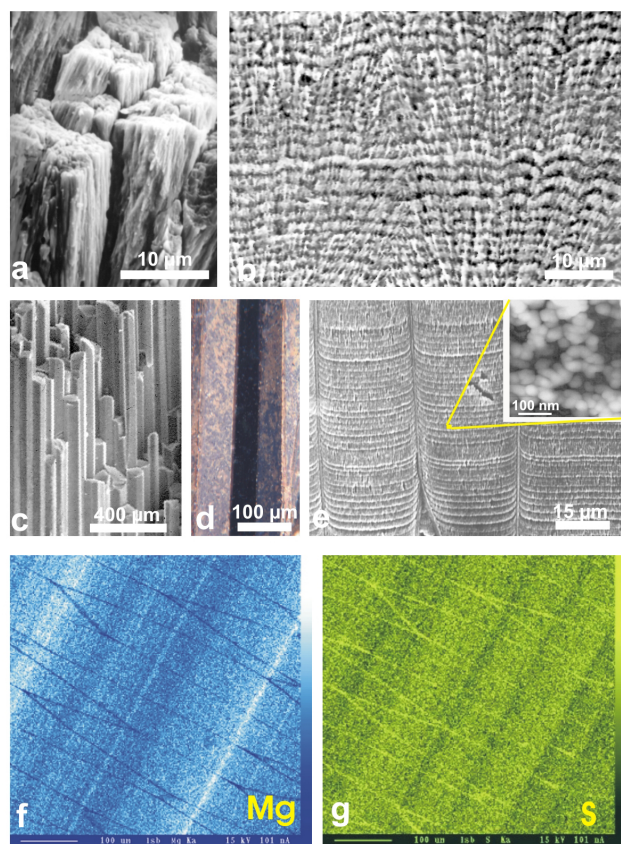
To create Ca-carbonate nanograins, calcium is actively transported through the cells of the basal ectoderm to the subectodermal space. Membrane sites involved and biochemical mechanisms have now been identified (Tambutté et al., 1995; Marshall, 1996; Zoccola et al., 2004). On the other hand, many analyses have shown that both glucidic and protein matrix components have compositional peculiarities favouring interaction with mineral ions. Glucids are sulfated and very acidic in composition (Dauphin, 2001), along with the aspartic/glutamic rich proteins (Mitterer, 1978; Constantz and Weiner, 1988). Taking into account the low isoelectric point of matrix components (the pH of the solution in which a given organic molecule is neutral), mineral ions certainly found numerous sites for fixing onto the organic framework (Clode and Marshall, 2003).

In this crystallization step, the polymeric organic framework may play a major role, being responsible for the position of the initial crystallization sites. Thus, after having been oriented by the underlying mineral surface, the organic framework itself may ensure the crystallographic coherence of the nanograins, an essential requirement for maintaining the overall crystallinity of fibres. From this standpoint, it is worthwhile to note that a close SEM examination of fibres (Fig. 14a) shows that the monocrystalline status of fibres is not so perfect than it could be expected if they were continuously growing purely mineral units, as proposed in the Bryan and Hill interpretation (Bryan and Hill, 1941).

Within the mineralizing layer, growth of mineral units progressively reduces the space available for the hydrated organic material secreted in the first step of the biomineralization cycle. At the end of this crystallization process, the organic phase simply appears as a cortex surrounding nanograins (see Fig. 10a–b).

##### 4.3.3 The coral fibre: a matrix-mediated biocrystal

In contrast to the still widely accepted monocrystalline concept, coral fibres belong to the category of matrix-mediated biominerals built by repeatedly produced micron-thick



**Fig. 14.** Similarity between the fine structure of coral fibrous tissue and a typical “matrix mediated” mollusc microstructure: the prismatic layer of *Pinna nobilis*. (a) Close view of coral fibres: each fibre is not a single crystal but a cluster of elongated subunits with similar polarizing behaviour (see Fig. 2). (b) Polished and etched surface in the fibrous zone of a coral skeleton: fibre subunits are still visible but the dominant pattern is the stepping growth mode. (c, d, e) Morphology, ultra-thin slide (polarized light) and etching surface in the *Pinna* prismatic shell layer. Calcitic prism morphology (c) is more precisely defined than coral fibres, due to their strong organic envelope. Monocrystalline behaviour (d) of the prisms is also better marked. However, an obvious similarity with the fine structures of coral fibres appears after etching (e). (f–g) Microprobe maps of a longitudinal section of the *Pinna* prismatic shell layer. Note the fine layering of minor element distribution, the high contrast between successive growth layers. Also remarkable is the independence in concentration changes between the two elements (Microprobe maps by C. T. Williams, NHM London).

growth layers. A comparison can be made between the coral fibre and the most classic of these polycyclic biocrystals: the calcitic prisms of *Pinna nobilis*, a pteriomorphid Pelecypod. The following points emphasize the similarity of the growth process between the molluscan prisms and coral fibres (Fig. 14).

- The micronic stepping growth mode (now proved also in fossil corals: Stolarski, 2003) is common to both

microstructures, with quite comparable patterns. The mean thickness of growth units and crystallographic co-ordination demonstrate a similar overall control.

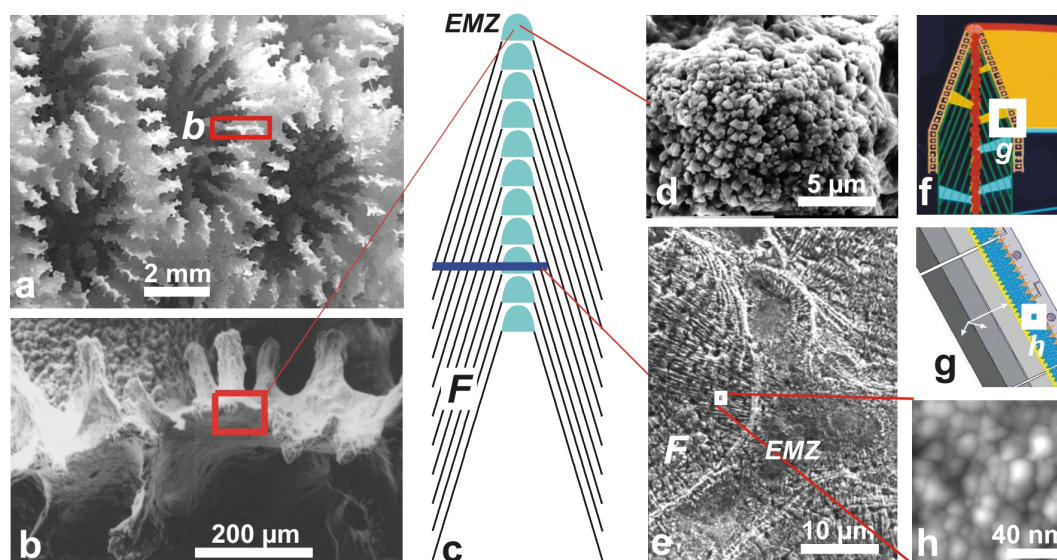
- In both cases growth layers are built by nanograins.
- From a biochemical standpoint, hydrated glycoconjugate components are present at a nanometre scale in both structures and exhibit the same spatial relationships (for XANES pictures of *Pinna* prisms see Dauphin et al., 2003).

#### 4.4 Relevance of the polycyclic model of coral fibre to recent fine scale chemical and isotopic measurements

During recent years, numerous studies have shown that proxies deriving from mineralized skeletons do not give us a very reliable information. Various discrepancies have been reported between environmental conditions and values measured on biological carbonates (Allison et al., 2001; Finch et al., 2003). When measurements are made using high resolution devices, chemical variability within a given specimen exceeds what can be explained by the influence of some physiological process, i.e. photosynthesis in reef corals (Rollion-Bard et al., 2003). Moreover, these unexpected variations also occur within deep sea corals (Blamart, 2002; Adkins et al., 2003). Obviously they are linked to the biomineralization process itself. Application of simple thermodynamic laws cannot provide us with an accurate interpretation of the environmental signal (Juillet-Leclerc, personal communication), and a new strategy (Lough, 2003) is now required.

To illustrate the rapid compositional changes that may occur in successive growth layers of polycyclic biominerals, comparison with the Pteriomorphid Pelecypod *Pinna nobilis* can be continued. In the *Pinna* prisms, microprobe maps of Mg and S (Fig. 14f–g) exemplify the variability in chemical composition between successive growth layers. The chemical layering strictly corresponds to the microstructural pattern and, moreover, concentration changes are different for the two mapped elements: sulfur varies independently of magnesium. Clearly, in contrast to common opinion, biological regulation of growth cannot prevent various influences to be recorded in the composition of growth layers. Although no equivalent document exists for coral skeletons, experiments have shown that mineralizing activity is very sensitive to physico-chemical conditions such as pH (Marubini et al., 2002).

The rapid changes in fractionation ratios evidenced by high resolution SIMS measurements in corals (Meibom, 2003; Rollion et al., 2003) are probably related to this high sensitivity, causing surprisingly rapid changes in environmental signals. As measurements made by usual sampling methods (even by computer driven hole driller, laser or ICP-MS) involve several growth layers (i.e. several environment



**Fig. 15.** From corallite morphology to skeletal nanograins of a septal growth layer in *Favia stelligera*. Measurements now possible at the micrometre range impose a good understanding of the skeletal features. (a–b) corallite (a) and septum growth edge (sight of the top); (c) scheme of superimposed growth layers (EMZ: early mineralization zones); (d) tiny isodiametric crystals in EMZ; (e) microstructure in the median part of a septum (*F*: fibre zone after etching); (f–g) scheme of growing surface with basal ectoderm (see also Fig. 13); (h) nanometre structure of a fibre growth layer.

recording units) variations between individual growth layers are probably still higher than reported in recent papers. Only the NanoSIMS method approaches the relevant measurement level.

Interestingly, the first mapping experiment applying the NanoSIMS method to coral fibrous tissues (Meibom, 2004) has produced a result that provides additional support to the hypothesis for a biological control on the crystallization process in corals. A layered Mg signal has been observed, repeatedly produced within each fibre growth layer, a result that allows suggesting a role for Mg in regulation of the crystallization process.

No doubt that until recently, the resolution of sampling methods was not adequate for the scale at which environmental information is recorded through biomineralization. Figure 15 summarizes the different structural level in coral skeletons. To develop the “new strategy” (Lough, 2003), this needs to be elaborated to improve the use of corals as environmental archives. A detailed understanding of the biomineralization process is a prerequisite. Precise relationships between environmental conditions and compositions in a given biomineral layer need to be established. Reciprocally, signals recorded within each skeletal growth layer acting as the Environment Recording Unit have to be measured at an appropriate scale.

## 5 Conclusions

1. Coral fibres are built by superimposition of a few micron-thick growth layers.
2. Growth layers are made of mineral nanograins densely packed within an organic component.
3. Concentric growth patterns visible within a given skeletal unit indicate that the growth process is coordinated at an overall scale by polyp physiology.
4. Due to cyclicity in the biomineralization process, each growth layer acts as an “Environment Recording Unit”.

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